PATENT SPECIFICATION

NO DRAWINGS

Inventor: EDWARD WALTON

1.187.824



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-C:2 C(1F3K4, 1E5K4, 3A10E4B3, 3A10E5E, 3A13A4A4, 3A13A4F1, 3A13A4F2, 3C:5A4, 3C:5C5, 3C:5E1, 3C:3E2, 214, 215, 22Y, 220, 25Y, 250, 252, 253, 28X, 30Y, 32Y, 323, 34Y, 342, 36Y, 360, 361, 362, 365, 366, 368, 45Y, 45X, 50Y, 503, 593, 598, 601, 603, 62X, 63X, 648, 652, 662, 668, 67X, 670, 680, 682, 173-198-289, 177-771-779, KK, KM, KY, LH) Index at acceptance:

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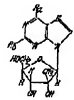
COMPLETE SPECIFICATION

Nucleosides and their Preparation

We, MERCE & CO. INC., a corporation duly organised and existing under the laws of the State of New Jersey, United States of America, of Rahway, New Jersey, United States of America, do bereby declare the Invention, for which we may that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:-

This invention relates to nucleosides.

The novel compounds of the present invention have the following structural formula:



In which each of R₀ and R₁₀, which may be the same or different, is a hydrogen or halogen amm or a hydroxy, C₁₋₁ alkyl, amino, C₁₋₀ alkylamino, di(C₂₋₀ alkyl)amino, mercapto or C₂₋₀ alkylinio radical.

Compounds of the present invention may be useful in the preparation of various 2'-methyl nucleotides, which may be useful in the study of nucleic acid metabolism, by their reaction with phosphorus compounds.

Typical values of R₀ and R₀, spart from those specifically mentioned above, are methyl, ethyl, propyl, methylamino, dimethylamino, ethylamino, diethylamino, propylamino, dipropylamino, chlorine, brumine, methylthic, methylthic and propylimino.

The compounds of the present invention are prepared in general by a two-step process. The first step in this process, Step A, is carried out by treating 2 2,3,5-tri-O-acyl-2-methyl-D-ribofuranosyl halden. 10

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with a chloromercuri 2,6-substituted purine of the formula:

in a solvent to form 9-(23,5-tri-O-acyl-2-methy!-D-ribofuranosyl)-2,6-solvationed purine limerine diates of the formula:

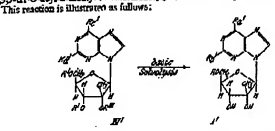
in which each of R, and R, is halosen, hydrogen, hydroxy, C, alkyl, acylamino or acyl C₁₋₃ alkylamino, each of R', R" and R", which may be the same or different, is an acyl group and X is a halogen atom. It is preferred that essentially stoiching the resemble of the resemble of the resemble of the resemble of the resemble. acyl C₁₋₃ anylamina, each of K. R. and K., which may be the sense a matchine is an acyl group and X is a halogen atom. It is preferred that resembly carried out in a metric amounts of the reactions be used. The reaction is preferably carried out in a temperature range of from 25°C to 150°C, particularly 100°C to 140°C, for a period of time sufficient to complete the reaction. This time is usually from 15 minutes to 5 hours; the higher the temperature, the quicker the reaction. The selection of the solhours; the higher the temperature, the quicker the reaction. The selection of the solhours; the higher the temperature, the quicker the reaction. The selection of the solhours; the higher than a long as it is an inert solvent and that it boils in a range of about 25°C to 150°C. Examples of such solvents are beazene, dibutyl ether, cyclo housen, toluene and kylene, preferably toluene and kylene.

The 2.55-tri-Q-acyl-2-nucthyl-D-ribofuranosyl halides used as starting materials are claimed in and may be prepared by processes described and claimed in the specification of our copending application No. 51812/69 (1,187,825). These processes comprise explaining 2-C-methyl-D-ribono-y-lactone to form 2,35-tri-Q-acyl-2-C-methyl-Q-sh-D-ribono-y-lactone, which is reduced with a dialkyl borane to produce 2,35-tri-Q-acyl-2-C-methyl-Q-sh-D-ribonoranose, which is further acylated to form 1,2,35-tri-Q-acyl-2-C-methyl-Q-sh-D-ribonoranose, which is further acylated to form 1,2,35-tri-Q-acyl-2-C-methyl-Q-sh-D-ribonoranose and converted into the ribofuranosyl halide by a halogenation repisoment reaction in a suitable colvent.

Those compounds of the present invention of Formula I', in which each of R. R. is a hydrogen or halogen arms or a hydroxy, C₁₋₅ alkyl, amino, C₁₋₂ alkylamino a

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where in Formula IV' each of R. and Ro is a hydrogen or halugen atom or hydroxy, C., alkyl, acylamino or acyl(C., alkyl)amino radical and in Fermula I' each of R. and R. is a hydroxen or halogen atom or a C., alkyl, hydroxy, amino or 30 C,- alkylamino radical.

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The solvolysis reaction is carried out in the presence of a basic catalyst in an appropriate solvent, preferably in a temperature range of from 5°C to 150°C, por ticularly 65°C to 90°C, in a reaction time of from 15 minutes to 5 hours. The length of reaction time is dependent upon the temperature, the catalyst and solvent used. Examples of basic catalysts are alkali metal and alkaline earth metal inorganic basis and their corresponding alkonides, solutions of ammonia, amines and substituted

and their corresponding automors, commons of summons, since sales and substitutes amines. Suitable solvents are $C_{i,-}$ alcoholz, preferably methanol.

In another aspect of the present invention, those compounds of Formula I', in which one or both of R_i , and R_b , is an amino, $C_{i,-}$ alkylamino or $di(C_{i,-}$ alkylamino radical, are prepared by an aminolytic reaction of those intermediate 9-(2,3,5-tri-O-ctyl 2-methyl. D-ribofuronoxyl.)-2,5-substituted purioes in which the 2 and/or 6 purioe position is substituted with a halogen, designated IV'',

The reaction is illustrated as follows:

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where at least one of $R_{c''}$ and $R_{d''}$ is a halogen atom, R in the aminolysis reagents is a C_{1-c} alkyl-radical and one of $R_{2''}$ and $R_{b''}$ in Founda I'' is amino, C_{0-c} alkylamino or di(C_{1-c} alkyl)amino.

The emmolysis reaction is carried out in the presence of ammonia, a C_{1-c} alkyl-amino or a di(C_{1-c} alkyl)amino, preferably at a temperature from 25°C to 150°C and preferably 85°C to 100°C in a reaction time of from 15 minutes to 5 hours. Examples of amines are methylamine, directlylamine, directlylamine, manufactors of amines are methylamine, dimethylamine, ethylamine, diethylamine, propylamine and dipropylamine.

In another expect of the present invention the compounds of Formula I'', in which one or both of R_1 —and R_2 —is or are message or C_{1-2} alkylthic are prepared by a mercaptolysis reaction, the preferred reagents being thiomea and professibly an formula MSR, where R is C_{1-2} alkyl and M is an equivalent of a mend, professibly an analysis of the professible of t alkali metal or an alkaline-carth metal, although any mercapnolysing agent capable of introducing a mercapio or C₁₋₁ alkylling group may he used. In the reaction scheme that follows, one or both of R_e—and R_e—in Formula IV" is a halogen atom, and each of R', R" and R" is anyl:

When the mercaprolysis reactant is thioures the anyl blocking groups R', R" and R" are not removed and the resulting intermediate must be subjected to having

and K''' are not removed and the resulting intermentary must be subjected to dear solvelysis in order to obtain the compounds of the present invention, Compound I'''.

The necropolysis reaction is carried out in the presence of thioures or a ment salt of a C₁₋₅ alkylthiol, preferably at a compensative of from 25°C to 50°C, particularly 65°C to 50°C, in a reaction time of from about 15 minutes to about 5 houre.

Preferred are the alkali metal and alkaling-earth metal salts of alkylthiols, e.g. sodium about 15 minutes and account metal and alkaling-earth metal salts of alkylthiols, e.g. sodium account of the compensation of the contraction metal-account metal-accoun methenethiolate, sodium ethanethiolate, sodium isopropanethiolate, potassium methansthiolore and colorum methanethiolate,

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4	the second of the present invention are	
	Representative of the novel compounds of the present invention are	
	9 (2-methyl-D-ribulusnosyl)-2-methylpurine	
	9-(2-mettyl-D-ribalmanosyl)-6-methylpurine 9-(2-mettyl-D-riladmanosyl)-2,6-dimethylpurine	5
_	9-(2-methyl-D-ribofuranosyl)-2-ethylpurine	3
5	9 (2-methyl-D-ringuranosyl)-6-ethylpurne	
	9 (2-methyl-D-rithfuranosyl)-2,6-diethylputine	
	o /2	
	o in maket. (1-infurishtsv1)-0-b(0py1pur) iii	10
44	o /> mathyl_/)atibofittanosy/)=2.0-dipropy/pumpo	•••
10	o 23 mather Darbonitzinosvi)-2-aminopurine	
	A / D material D-randfur 200 EVI) - O-MILLING DUTING	
	0. /2e/w/_/1_ribofnr8706V/)-2.0=181mmopuuse	
	ο /2_methol_/)_ribofuranosv()-2-iiicinyiamuropur με	15
15	C >>	
_	9-(2 methyl-D-ribofitranosy!)-2,6-dimethylaminopuring	
	9-(2-methyl-1)-ribofurancsyl)-2-ruhylaminopurine	
	9-(2-methyl-D-ribofurancyl)-6-chylaminopurine	
	9-(2-methyl-D-cibofuranosyl)-2,6-dicthylaminopurine	20
20	9-(2-methyl-D-riboturanoxyl)-2-hydroxypurine 9-(2-methyl-D-riboturanoxyl)-6-hydroxypurine	
	9-(2-methyl-D ribofursnosyl)-2,6-dihydroxyputine	
	A /a	
	A /A analysis II as had neutrocky leverticity in the investment of	25
25	A A .g., y Th _!Laften Accol., 7_merby/group)=0*HCU1VAUU-U-	2,2
20	o /o-merkyLD -chocuranosyl)-2-amino-o-inculyianimopulane	
	Δ./9_eeekv)./ λ.whotuf100sV1.~/•IIICIDY1-V-IIVIVATPMM*	
	A /A A-Bast D Bofters moses to A-months v-0-months und	
	o /a	30
30	9-(2-methyl-D riboturanosyl-7-hydroxy-G-aminopurine	
	9-(2-monyl-D-ribofuranosyl)-7-methylamius-6-hydroxypurine	
	9-(2-methyl-D-ribofuranosyl)-2-hydroxy-6-methylaminopurine	
	9-(2-methyl-D-ribofuennosyl)-2-dimerhylaminopurine	
	9-(2-methyl-D-ribofuranosyl)-6-dimerhylaminopurine 9-(2-methyl-D-ribofuranosyl)-2-methylamino-6-dimethylaminopurine	35
35	9-(2-methyl-D-modratosyl) 2-mercapropurine	
	9-(2-methyl-D-ribofuranosyl)-6-mercaptopurine	
	9-(2-methyl-D-ribofuranosyl) 2.6-dimercaptomine	
	o //	40
40		44.7
40	o (2-mail-dLTh-ribAfirentesti) 2-mercapin-o-intrinyindal-datoy-umas	
	9.72-methyl-D-ribefuranceyl)-2,6-dicbloropume	
	9-(2-methyl-D-ribofurancoyl) 2,6-dichloropurine 9-(2-methyl-D-ribofurancoyl) 2-chloropurine	
	0_/7_methyl=/)-rthcft://2703V1}-2-0rth1098time	45
45	0 /2 hard-Marshoth rennated before the compositions	
	9_/?auterbyle/)-cthofurandsyl)-6-chloropurine	
	9-(2-methyl-D-ribofuranciyl)-2,6-dibromopurine	
	and the same and house	
	Compounds of the present invention have a variety of valuable uses and have	
		50
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	growth of KB cells has been shown to be markedly suppressed by compounds of the	
	present invention as is the incorporation of hypoxambine into acid insoluble RNA. Compounds of the present invention are therefore useful as animetabolities, as cell Compounds of the present invention are therefore useful as animetabolities, as cell	
	possess favourable cytotoxicity characteristics considered with their cell growth depres-	55
55		
	Compounds of the present invention may also be converted to nucleotides by	
	such, they are useful in a formulation of media for study of nucleic acid metabolism, cells. These nucleotides may also be useful in the study of nucleic acid metabolism.	60
60	The following examples illustrate the compounds of the present invention.	
	Examples 1 and 1A being comples of the invention claimed in the specification of	
	Examples I and IA being complete of the inventor Complete in the speciments	

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5 1,187,824 our coperading application No. 51812/69 (1,187,825). In the Examples all parts are by weight and the word 'Dower' is a trade mark. ETAMPLE 1 Preparation of 2,3,5-Tri-O-benzoyl-2-methyl-D-ribofuranosyi chloride This example shows the synthesis of a novel starting material used in the preparation of the compound of the present invention. A solution of 5 g. (30.8 millimoles) of 2-C methyl D ribono-y-lactone in 100 ml. of dry pyridine is cooled to about 5 °C, and treated with 17 ml. of benzoyl chloride. The mixture is heated to 65--/0°C, for 4 hours and kept at room temperature for 16 hours. The reaction mixture is stirred with 2 ml. of water for 25 minutes to de-10 10 compose unreacted benzoyl chloride, and the pyridine is removed at reduced pressure. The thick residue is dissolved in 100 mL of chloroform and the chloroform solusure. The Linex residue is discoved in 100 ml. of chloroform and the chloroform solution is washed with three 50-ml, portions of 10 percent hydrochloric acid, two 50-ml, portions of 10 percent sodium blearbonate and two 50-ml, portions of water. The dried chloroform solution is concentrated and the residue is dissolved in other. The ethereal solution is concentrated to 250 ml, and after being cooled to 5°C, for several hours gives 8.8 g. (60%) of 2,3,5-tri-O-benzovi-2-C-methyl-D-ribo-y-lactone, m.p. 138—140°C. 15 15 A solution of 7 g. (14.7 mmole) of 2,3,5-tri-O-benzoyl-2-C-methyl-D-ribonoγ lactore in 30 ml. of dry tetrahydrofuran is cooled, under nitrogen, and treated with
58.3 ml. of 1 M discemdaryisoamylborane. The reaction solution is kept at roun temperature for 16 hours. After the careful addition of 6 ml, of water, the mixture is refluxed for 0.5 hour. The mixture is cooled to about 5°C, and 11.5 ml. of 30% hydrogen peroxide is added while the pH is maintained between 7 and 8 through die additition of about 7 ml. of 3N sodium hydroxide. The mixture is extracted with six 50ml. portions of chloroform and the extracts are washed with several purtions of water.
The ramaining peroxides are removed by washing the chloroform solution with 90%
forrous sulfate. Concentration of the chloroform layer gives 7.8 g. of crude product
as a syrup. The product is purified by chromatography on 200 g. of silica gel in a
mixture of benzene and cityl accrate (4:1). From the column, fractions are obtained
which contain 2.5 g. (37%) of 2,3,5-tri-O-benzoyl-2-C-methyl-(α and β)-D-ribnfurancese. 20 75 30 A solution of 4.2 g. (8.8 millimoles) of 2.3.5-tri-O-benzoyl-2-C-methyl-(α,β)-D-ribofurances (containing a small amount of 3.5-di-O-benzoyl-2-C-methyl-(α,β)-D-ribofurances) in 80 ml. of dry pyridine is cooled and treated with 8.0 ml. (68 millimoles) of benzoyl chloride. The mixture is heated at 90°C, for 4 bours and cooled to 95 35 about 50C. A small amount of water is added and the unxure is stirred for 0.5 hour to decompose excess of benzoyl chluride. The reaction mixture is concentrated and the residue is dissolved in chloroform. The chloroform solution is washed with three the residue is dissolved in Chia otomi. The character standard is wasness with three 50-ml. partients of 10% hydrochloric acid, three 50-ml. partients of saturated sodium bicarbonane and three 50-ml. portions of water. The third chloroform layer is concentrated to 5.1 g. of an oil. Addition of 50 ml. of other gives 2.16 g. (47%) of 12.3 5-tetra-0-benzoyl-2-C-methyl-B-D-ribolurances, m.p. 155-156°C. Concentration of the filtrate gives 2.9 g. (57%) of essentially pure 1,23,5-tetra-0-benzoyl-2-C-methyl-D-ribolurances as an oil. a-D-ribofurance as an oil.

To 100 ml. of a saturated solution of hydrogen chloride in other is added 2 ml. of accept chloride and 1.5 g. (2.6 milliandes) of 1,23,5-ners-O-hentopt-2-C-methyl-45 45 B-D-ribofurance. The solution is kept at room temperature for 2 hours and the effect is removed at reduced pressure. Five 21-ml. partions of dry minene are successively removed at reduced pressure from the residue. The residue is dissolved in dry ether and quickly washed with cold saturated addition bicarbocate and tinally with cold water. After being dried the ethereal solution is communicated and a residue of 2,3,5-tin Department of the design of the design of 2,3,5-tin Department of the design of t 50 50 tri-O-benzoyl-2-C-methyl-B-D-ribuluranusyl chloride is obtained. EXAMPLE 1A Preparation of 2,3,5-Tri-O-benzoyl-Z-methyl-D-rihofurnosyl Bromide 55 Preparation of 2,3,5-Tri-O-benzoyl-Z-methyl-D-riboturenosyl Bromide
A solution of 1.5 g. (2.6 millimates) of 1,2,3,5-terra-O-benzoyl-Z-C-methyl-a-Driboturenose as prepared in Example 1 in 7.3 ml. of sectic acid is treated with 0.25
ml. of sectyl bromide and 7.5 ml. of a 32% (w/w) solution of hydrogen bromide in
acetic acid. The mixture is kept at 25°C, for 24 hours. The mixture is concentrated
and a portion of day toluran is distilled, at reduced pressure, from the residue to
remove excess hydrogen bromide and acretic acid. The residue is dissolved in dry
other and quickly wasted with oald acrusted sodium bicarbonate and finally with
cold water. After being dried the ethereal solution is enceenbrated and a residue of 55 cold water. After being dried the enternal solution is concentrated and a residue of 2,3,5 cm-O-benzuyl-2-C-methyl-\$-D-ribemunesyl bromide is obtained.

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1,187,824 EXAMPLE 2 Preparation of 9-(2,3,5-tri-O-benzuyl-2-methyl-D ribofuranosyl)-2-acetamidu-6 Preparation of 9-(2,3,5-tri-O-benzuyl-2-methyl-D ribofuranosyl)-2-aceramido-6-hydroxypurine

About 25 ml. of xylene is distilled from a suspension of 5.95 grams (0.014 mole) of chloromerami 3-acetamido-6-hydroxypurine in 175 ml. of xylene to remore the last tracts of water. The suspension is cooked to 25°C, and 2,3,5-uri-O-benzoyl-2-methyl-D-ribofuranosyl chloride prepared from 8.1 grams (0.014 mole) of 1,2,3,5-uri-O-benzoyl-2 methyl-D-ribofuranose in 25 ml. of dry xylene is added. The nursure is stirred and heated at a temperature of from about 50°C, to about 100°C. The solid changes from a granular form to florundent. After being refluxed for one bour, the list incure is filtered, which removes the undissolved solids. Leaching the solids with three 50-ml. portions of boiling chloroform removes additional soluble nursure. 10 with three 50-ml, portions of boiling chloroform removes additional soluble product and leaves insoluble starting chloromercuri derivatives and inorganic sales. The original filtrate is diluted with two volumes of perroleum other and the solid which separates is dissolved in the chloroform solution obvained above. The chloroform solution plus 15 an additional 100 ml. is washed with two 75 ml. portions of 30% potassium lodde solution and two 75 ml. portions of water. The dry chloroform layer is concentrated and 9-(2,3,5-tri-O-benzoyl 2-methyl-D-ribathranesyl)-2-amin-6-hydroxypurine is ob-15 EXAMPLE 3 Freparation of 9 (2,3,5-tri-O-henzayl-2-methyl-D-ribofuranosyl)-6-N-methylbenzamidupusinc 20 About 150 ml. of sylene is distilled from a suspension of 9.5 grams (19.5 millimoles) of chlorencercuri 6-N-methylbenzamidupurine in 500 ml. of xylene. The mixmoies) of coloremercian or templocoles management in 300 m. of ayear. The most use is cooled and a solution of 2,3,5-tri-O-benzoyl-2-methyl-D-ribofuranosyl chloride (from 8.2 grams [14.1 millimoles] of 1,2,3,5-texts-O-benzoyl-2-methyl-D-ribofuranose) in 50 ml. of dry zylene is added. The reaction mixture is stirred and refluxed for 30 minutes. The hot mixture is filtered and 3 grams of unreacted stationary delegations are refluxed to decrease and the 25 25 ing chloromercus; purine is recovered. The fittrate is tunceturated to dryness and the residual oil in 300 ml. of chlorotem is washed with two 80-ml. portions of 30% potassium iodide solution and two 80-ml. portions of water. The residual oil obtained an. after removal of the chloroform is chromatographed on a short column of 140 grams of acid-washed alumina in 9 to 1 benzene-chloroform. Fractions are combined and consumated giving 9-(2,3,5-tri-O-benzoyl-2-methyl-D-ribofuranosyl)-6-N methylberoxmidopurioc. 35 EYAMPLE 4 Preparation of 9-(2,3,5-tri-O-benzoy)-2-methyl-D-ribofuranosyl)-6-chloroparine Proparation of 9-(2,3,5-tri-O-benzoyl-2-methyl-D-ribofuranosyl)-6-chloroparine About 100 ml. of xylene is distilled from a autpension of 6.55 grams (16.8 millimoles) of chluromercuri-6-chloroparine in 460 ml. of xylene in order to remove the last traces of water. A solution of 9.05 grams (16.8 millimoles) of 2,3,5-tri-O-benzoyl-2-methyl-D-ribofuranosyl bromide in 40 ml. of dry xylene is added to the stirred suspension at 25°C. The mixture is refluxed for 2 hours. The hot mixture is filtered to remove insulvide material. The filtrate is concentrated to 150 ml. and diluted with 300 ml. of petroleum ether. The mixture is kept at 5°C. for one hour and filtered. The solid is washed with three 20 ml. portions of petroleum ether and dried. The crude product is dissolved in 300 ml. of hot chloroform and washed with two 80-ml. portions of 30% potassium iodida solution and two 80-ml. portions of water. The dried (Mg5O.) chloroform layer is concentrated, and 9-(2,3,5-tri-O-benzoyl-2-methyl-D-ribofuranosyl)-6-chloropurine is obtained. The product is purified by chromato-45 D-rihoturanoxyl)-6-chloropurine is obtained. The profiner is purified by chromatography on a short alumina column in chloroform. 50 Preparation of 9-(2,3,5 tri-U-benzoyl-Z-methyl-I)-ribofuranosyl)-2,6-dibinzanoidopurine

About 100 ml. of xylene is distilled from suspension of 5.01 grams (8.43 millimoles) of diboromercuri, 2,6-dibenzamido purine in 370 ml. of xylene to remove last traces of water. The suspension is cooled to room temperature in a solution of 4.55 grams (8.43 millimoles) of 2,3,5-tri-U-benzoyl-Z-methyl-D-rimfuraturyl bromide in 37 ml. of dry xylene is added while the suspension is being seitred. The misture is refused for 2 hours and filtered hot which removes insulable material. The filtrate is dillined with 400 ml. of petroleum other and cooled in an low bath. The solid is removed and dried. The product is obtained as a complex with mercuric halide. The HEALIPLE S 55 55 60 moved and dried. The product is obtained as a complex with mercuric halide. The product is dissolved in 100 ml, of chloroform and washed with two 40-ml, portions

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7	1,187,824	7
	of 30% potassium iodide solution and two 40-ml, partions of water. The dried (MgSO ₂) chluruform solution is concentrated at reduced pressure to give 9-(2,3,5-tri-O-benzoyl-2-methyl-D-riboforanosyl-2,5-dibenzamidoporine.	
5	EXAMPLE 6 Preparation of 9-(2,3,5-tri-O-benzoyl-2-methyl-D-ribofurancsyl)-6-methyl-purine A suspension of 3.7 grams (10 millimoles) of chloromercuri 6-methyl-purine [Davoll and Lowy, J. Am. Chem. Soc. 73 1650 (1951)] in 200 ml. of sylene is dried by distilling about 50 ml. of sylene. The cooled suspension is treated with 4,94 grame (10 millimoles) of 7.3.5-tri-O-benzoul-2-methyl-purine in the cooled system of the cooled syst	5
10	(in millimates) of 2,3,5-ri-D-benzoyl-2-methyl-D-ribofursmosyl chloride dissolved in 30 ml. of dry xylene. The mixture is stirred and refluxed for 2 hours and it is filtered to remove insoluble material. The filtrate is chluted with 4 volumes of petrolcum other and, after being enoted for about 2 hours in an ice bath, the mixture is filtered. The solid is dissolved in 200 ml. of chloroform and weeked with two 30-ml. pertions of 20% aqueous potassium iodide solution. The chloroform layer is dried (anhydrous MgSO ₂) and concentrated to a residue of amorphous 9-(2,3,5-ri-O-benzoyl-2-methyl-	10 15
_	ribraturanosyi)-5-methylpurine,	D
	Frample 7 Preparation of 9-(2,3,5-tri-O benzoyl-2-methyl-D-zibofuranosyl)-6- benzamidopurine	
20	A suspension of 2.92 grams (5.95 millimoles) of finely ground chloromercuri 6-betramidopurme in 200 ml. of sylene is dried by distilling 100 ml. of sylene. The mixture is cooled and a solution of 23,5-tri-O-benzoyl-2-methyl-D-ribofuranosyl chloride funde from 3.45 grams (5.95 millimoles) of 1.23,5-tetra-O-benzoyl-2-methyl-D-ribofuranosyl m 30 ml. of dry sylene is added. The mixture is client and added.	20
25	for 80 minutes. The hot mixture is filtered and the solid is washed with 25 ml. of hot sylane. The filtrate and washings are diluted with 400 ml. of petroleum ether, and after being kept at 5 °C. for 20 hours, the mixture is filtered. The solid is dissolved in 300 ml. of chlerederm and the solution is washed with two 20-ml, portions of 90% potential modifies solution and two 20 ml. portions of water. Concentration of the	25
30	dried chloroform layer gives amorphous product which is chromatographed on 70 grams of alumina in othyl acetate chloroform (I:4). Fractions showing only one zone (Re 0.65) after thin layer chromatography on alumina in othyl acetate-chloroform (1:4) are combined and concentration of the solvent gives 9-(2.3,5-tri-O-benzuyl-2-methyl-D-ribofuranceyl)-6-benzamidopurine as an amorphous solid.	30
35	Example 8 Preparation of 9-(2-Methyl-D-nibofuranosyl)-6-dimethylaminopurine A suspension of 1.0 gram (1.57 millimale) of 9-(2,3,5-ml-O-henzoyl-2-methyl-D-zibofuranosyl)-6-chloropucine as prepared in Example 4 in 25 ml. of methonol continues of Sylvanosy	35
40	mining 6.5 grams of directly langue is heated for 10 hours in a sealed nube at 100°C. The solution is concentrated at reduced pressure and the residue is dissolved in 25 rol. of water. The water solution is washed with five 8-nd. purrious of benzene and then treated with 2 grams of Dower II-X8 which is a strongly basic anium-exchange resin having a stryrene divinyl benzene polymer matrix and containing quaternary	40
45	ammonium groups. It has an average particle size in the range of 50—100 mesh. It is manufactured by the Dow Chemical Co. of Malland, Michigan (see Page 1576, 7th Md., Merck Index, Merck & Co., Inc., Rahway, N.J. The resin is filtered and washed with three 25 ml. pertions of water. The filtrate is enterparated to dryness and 9-(2 methyl-D sibofurmosyl)-6-dimethylaminoparine is obtained.	45
50	Example 9 Preparation of 9-(2-methyl-D-ributuransyl)-2,6-diaminopurina A mixture of 1.2 grams (1.37 millimules) of 9-(2,3,5-tri-O-benzoyl-2-methyl-D-ributuranosyl)-2,6-dibutaminologurine as prepared in Rrample 5 in 12 ml. of dry methanol is treated with a solution of 97 mg. of (4.2 millimules) of sodium in 12 ml.	50
55	of methanol. The mixture is refluxed for 3 hours and the resultant solution is concentrated at reduced pressure. The residue is dissolved in 24 ml. of water and the pH is adjusted to about 6.5. The aqueous solution is extracted with five 10-ml. portions of chloroform to remove methyl hencestre and concentrated at reduced pressure to a recidue containing 9-(2-methyl-D-ribufuranceyf)-2,6-disminopurine.	55
60	Example 10 Preparation of 9-(2-methyl-D-ribothyranceyl)-purine-6-thiol	60

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in which each of R₀ and R₁₀, which are the same as or different from one another, is a hydrogen or halogen atom or a hydroxy, C₁₋₃ alkyl, amino, C₁₋₅ alkylamino, dl(C₁₋₆ alkylamino, mercapto or C₁₋₆ alkylamino radical.

2. 2'-Methyladenosine.

3. 9 (2-Methyl-D-ribufuranosyl)gusnine.

4.9 (2-Methyl-D-ribufuranosyl)-purine-6 thiol.

5. The process that comprises, in a first step, treating a compound of the formula:

15

in which K', R'' and R''' are the same or different acyl groups and X is a halogen atom with a compound having the formula:

in which each of R, and R₀, which are the same as or different from one another, is a halogen or hydrogen atom or a hydroxy, C₁₋₄ alkyl, acytamino or acyt-(C₂₋₄ alkyl)-amino radical, in a solvent to form a compound of the formula:

15



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